

General

Guideline Title

ACR Appropriateness Criteria® pediatric Hodgkin lymphoma.

Bibliographic Source(s)

Terezakis SA, Metzger ML, Constine LS, Hodgson DC, Schwartz CL, Advani R, Flowers CR, Hoppe BS, Ng A, Roberts KB, Shapiro R, Wilder RB, Yunes MJ, Expert Panel on Radiation Oncology-Lymphoma. ACR Appropriateness Criteria® pediatric Hodgkin lymphoma. [online publication]. Reston (VA): American College of Radiology (ACR); 2012. 16 p. [54 references]

Guideline Status

This is the current release of the guideline.

This guideline updates a previous version: Constine LS, Hoppe RT, Colman M, Deming RL, Mendenhall NP, Morris DE, Ng A, Wolkov HB, Yahalom J, Chauvenet AM, Hudson MM, Winter JN, Mauch PM, Expert Panel on Radiation Oncology-Hodgkin's Disease Work Group. Pediatric Hodgkin's disease. [online publication]. Reston (VA): American College of Radiology (ACR); 2006. 30 p.

Recommendations

Major Recommendations

ACR Appropriateness Criteria®

Clinical Condition: Pediatric Hodgkin Lymphoma

<u>Variant 1</u>: 12-year-old girl with CS IIA NSHL, three sites including nonbulky mediastinal and bilateral neck disease with rapid early response after initial chemotherapy and complete response at the end of chemotherapy.

Treatment	Rating	Comments	
Chemotherapy with or without Radiation Therapy			
4 cycles ABVD alone	6	Data for ABVD are sparse for pediatric patients.	
6 cycles ABVD alone	3	This is typical therapy in adults, but probably more therapy than necessary for children.	
2 cycles ABVD + 20-26 Gy IFRT	5		
Rayings Stelle D1;2230 Usually I filt fapprop	riate; 4,5,6 May be ap	propriate; 7,8,9 Usually appropriate	

6 cycles ABV Dre 2002 ot Gy IFRT	Rating	Comments		
4 cycles ABVE-PC alone	6			
3-4 cycles ABVE-PC + 20-26 Gy IFRT	5	More therapy than is necessary.		
5 cycles ABVE-PC + 20-26 Gy IFRT	3			
2 cycles O(E/P)PA alone	8			
2 cycles O(E/P)PA + 20-26 Gy IFRT	5			
2 cycles O(E/P)PA +>26 Gy IFRT	3			
2 cycles O(E/P)PA + 2 COP(P/Dac) + 20-35 Gy IFRT	2			
2 cycles O(E/P)PA + 4 COP(P/Dac) + 20-35 Gy IFRT	1			
4 cycles VAMP alone	3	Should receive consolidated radiation.		
4 cycles VAMP + 15-20 Gy IFRT	7			
4 cycles VAMP + 21-26 Gy IFRT	4	Too much radiation.		
2 cycles DBVE + 20-26 Gy IFRT	6			
4 cycles DBVE alone	3			
4 cycles DBVE + 20-26 Gy IFRT	6			
4 cycles COPP/ABV hybrid alone	6			
6 cycles COPP/ABV hybrid alone	4			
4 cycles COPP/ABV hybrid + 20-26 Gy IFRT	6			
4 cycles BEACOPP + 4 COPP/ABV	2			
4 cycles BEACOPP + 2 ABVD + 20- 26 Gy IFRT	2			
8 cycles BEACOPP + 20-26 Gy IFRT	1			
Rating Scale: 1,2,3 Usually not appropriate; 4,5,6 May be appropriate; 7,8,9 Usually appropriate				

<u>Variant 2</u>: 6-year-old girl with CS IIB NSHL with bulky mediastinal disease with rapid early response after initial chemotherapy and complete response at completion of chemotherapy.

Treatment	Rating	Comments		
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4 cycles ABV Declarate nt	Rating	Comments
6 cycles ABVD alone	4	
2 cycles ABVD + 20-26 Gy IFRT	3	
4 cycles ABVD + 20-26 Gy IFRT	6	
6 cycles ABVD + 20-26 Gy IFRT	4	
4 cycles ABVE-PC alone	7	Based on data from recently completed COG study. Data still under analysis.
3-4 cycles ABVE-PC + 20-26 Gy IFRT	8	
5 cycles ABVE-PC + 20-26 Gy IFRT	6	
2 cycles O(E/P)PA alone	1	
2 cycles O(E/P)PA + 20-26 Gy IFRT	3	
2 cycles O(E/P)PA +>26 Gy IFRT	3	
2 cycles O(E/P)PA + 2 COP(P/Dac) + 20-35 Gy IFRT	7	
2 cycles O(E/P)PA + 4 COP(P/Dac) + 20-35 Gy IFRT	5	
4 cycles VAMP alone	3	
4 cycles VAMP + 15-20 Gy IFRT	3	
4 cycles VAMP + 21-26 Gy IFRT	3	
2 cycles DBVE + 20-26 Gy IFRT	2	
4 cycles DBVE alone	2	
4 cycles DBVE + 20-26 Gy IFRT	3	
4 cycles COPP/ABV hybrid alone	4	
6 cycles COPP/ABV hybrid alone	5	
4 cycles COPP/ABV hybrid + 20-26 Gy IFRT	5	
4 cycles BEACOPP + 4 COPP/ABV	4	
4 cycles BEACOPP + 2 ABVD + 20- 26 Gy IFRT	4	
8 cycles BEACOPP + 20-26 Gy IFRT	2	
Rating Scale: 1,2,3 Usually not appropr	iate; 4,5,6 May be appro	opriate; 7,8,9 Usually appropriate

<u>Variant 3</u>: 16-year-old boy with CS IIIA (neck, mediastinum, para-aortic) nonbulky NSHL with slow early response after initial chemotherapy and complete response at completion of chemotherapy.

Treatment	Rating	Comments
Chemotherapy with or without Radiatio	n Therapy	
4 cycles ABVD alone	3	
6 cycles ABVD alone	4	
2 cycles ABVD + 20-26 Gy IFRT	3	
4 cycles ABVD + 20-26 Gy IFRT	5	
6 cycles ABVD + 20-26 Gy IFRT	4	
4 cycles ABVE-PC alone	3	
3-4 cycles ABVE-PC + 20-26 Gy IFRT	8	
5 cycles ABVE-PC + 20-26 Gy IFRT	8	
2 cycles O(E/P)PA alone	1	
2 cycles O(E/P)PA + 20-26 Gy IFRT	2	
2 cycles O(E/P)PA +>26 Gy IFRT	2	
2 cycles O(E/P)PA + 2 COP(P/Dac) + 20-35 Gy IFRT	8	
2 cycles O(E/P)PA + 4 COP(P/Dac) + 20-35 Gy IFRT	4	
4 cycles VAMP alone	2	
4 cycles VAMP + 15-20 Gy IFRT	3	
4 cycles VAMP + 21-26 Gy IFRT	3	
2 cycles DBVE + 20-26 Gy IFRT	2	
4 cycles DBVE alone	3	
4 cycles DBVE + 20-26 Gy IFRT	4	
4 cycles COPP/ABV hybrid alone	3	
6 cycles COPP/ABV hybrid alone	5	
4 cycles COPP/ABV hybrid + 20-26 Gy IFRT	4	
Ratings State CO23 Usually Photophypropr	iate; 4,5,6 May be appro	priate; 7,8,9 Usually appropriate

4 cycles BEACOPP + 2 ABVD + 20-	Rating	Comments	
26 Gy IFRT			
8 cycles BEACOPP + 20-26 Gy IFRT	3		
Rating Scale: 1,2,3 Usually not appropriate; 4,5,6 May be appropriate; 7,8,9 Usually appropriate			

<u>Variant 4</u>: 14-year-old girl with CS IIIB (supraclavicular, mediastinum, para-aortic + splenomegaly) MCHL with partial response at completion of chemotherapy.

Treatment	Rating	Comments
Chemotherapy with or without Radiation	Therapy	
4 cycles ABVD alone	2	
6 cycles ABVD alone	3	
2 cycles ABVD + 20-26 Gy IFRT	2	
4 cycles ABVD + 20-26 Gy IFRT	3	
6 cycles ABVD + 20-26 Gy IFRT	7	
4 cycles ABVE-PC alone	3	
3-4 cycles ABVE-PC + 20-26 Gy IFRT	4	
5 cycles ABVE-PC + 20-26 Gy IFRT	8	
2 cycles O(E/P)PA alone	1	
2 cycles O(E/P)PA + 20-26 Gy IFRT	2	
2 cycles O(E/P)PA +>26 Gy IFRT	2	
2 cycles O(E/P)PA + 2 COP(P/Dac) + 20-35 Gy IFRT	3	
2 cycles O(E/P)PA + 4 COP(P/Dac) + 20-35 Gy IFRT	8	
4 cycles VAMP alone	2	
4 cycles VAMP + 15-20 Gy IFRT	2	
4 cycles VAMP + 21-26 Gy IFRT	2	
2 cycles DBVE + 20-26 Gy IFRT	2	
4 cycles DBVE alone	2	
4 cycles DBVE + 20-26 Gy IFRT	3	

4 cycles COPIT/ABX/harid alone	Rating	Comments	
6 cycles COPP/ABV hybrid alone	3		
4 cycles COPP/ABV hybrid + 20-26 Gy IFRT	5		
4 cycles BEACOPP + 4 COPP/ABV	6		
4 cycles BEACOPP + 2 ABVD + 20- 26 Gy IFRT	5		
8 cycles BEACOPP + 20-26 Gy IFRT	7		
Rating Scale: 1,2,3 Usually not appropriate; 4,5,6 May be appropriate; 7,8,9 Usually appropriate			

 $\underline{\text{Variant 5}}\text{: 7-year-old boy with CS IIA nLPHL isolated to the right iliac and inguinal nodes.}$

Treatment	Rating	Comments
Surgery alone	3	
Radiation Therapy Alone		
15-25 Gy IFRT	3	Inadequate data.
>25 Gy IFRT	3	Not recommended in this age group due to long-term side effects to the growing skeleton.
Chemotherapy with or without Radiatio	n Therapy	
3 cycles AVPC alone	7	Data from recent COG study are promising.
3 cycles CVP alone	7	
4 cycles ABVD alone	5	
6 cycles ABVD alone	3	
2 cycles ABVD + 20-26 Gy IFRT	6	
4 cycles ABVD + 20-26 Gy IFRT	4	
6 cycles ABVD + 20-26 Gy IFRT	2	
4 cycles ABVE-PC alone	4	
3-4 cycles ABVE-PC + 20-26 Gy IFRT	3	
5 cycles ABVE-PC + 20-26 Gy IFRT	2	
2 cycles O(E/P)PA alone	7	Assuming CR.
2 cycles O(E/P)PA + 20-26 Gy IFRT	6	
2 cycles O(E/P)PA+>26 Gy IFRT	3	Too much radiation.

2 cycles O(E/PDPPathPACOP(P/Dac) +	Rating	Overly toxic, unless newer techniques ridse IMRT are considered.
20-35 Gy IFRT		Currently there are no data on these newer techniques.
2 cycles O(E/P)PA + 4 COP(P/Dac) + 20-35 Gy IFRT	1	
4 cycles VAMP alone	4	
4 cycles VAMP + 15-20 Gy IFRT	7	
4 cycles VAMP + 21-26 Gy IFRT	6	
2 cycles DBVE + 20-26 Gy IFRT	6	
4 cycles DBVE alone	4	Inadequate data.
4 cycles DBVE + 20-26 Gy IFRT	5	
4 cycles COPP/ABV hybrid alone	4	
6 cycles COPP/ABV hybrid alone	3	
4 cycles COPP/ABV hybrid + 20-26 Gy IFRT	3	

Summary of Literature Review

Introduction

Historically children and adults were treated with the same chemotherapy regimens, radiation therapy fields, and doses. However, irradiation techniques suitable for adults produced significant morbidities in children, such as impaired musculoskeletal development, an increased risk for subsequent benign and malignant neoplasms, and cardiopulmonary toxicities that became unacceptable. A desire to reduce these morbidities has motivated the development of new treatment strategies for pediatric Hodgkin lymphoma (HL).

Epidemiology

Childhood HL comprises 6% of childhood cancers and is epidemiologically distinct from adult HL. A striking male/female predominance is found among young children (ratio of 4:1 for 3- to 7-year-olds, and 3:1 for 7- to 9-year-olds) whereas the ratio for older children is closer to that of adults (1.3:1). The disease is uncommon before age 5, and among children is most common in adolescence.

Evidence for a genetic predisposition exists and is relevant when counseling families. Siblings have a 2- to 5-fold increased incidence, and this rises 9-fold in same-sex siblings. Parent-child associations are reported. One study reported a 99-fold increased risk in monozygotic twins of patients, but no increased risk in dizygotic twins. The role of Epstein-Barr virus (EBV) in the pathogenesis of HL is well established. In one report, EBV early RNA1 was expressed in R-S cells in 58% of childhood cases. Of particular interest is that expression was age dependent — 75% of children under age 10 compared with 20% of older children. In addition, a history of infectious mononucleosis increases the risk for HL, and anti-EBV titers are elevated prior to diagnosis of HL.

Clinical Presentation

HL typically presents with a dominant nodal mass, with 90% of patients demonstrating contiguous lymphatic spread. Most children are diagnosed on the basis of supradiaphragmatic lymph nodes, with painless cervical adenopathy in 80% of cases. Mediastinal involvement occurs in 76% of adolescents, but in only 33% of 1- to 10- year-olds. About one-third of patients will have systemic "B" symptoms of fever (temperature >38°C), drenching night sweats, and/or unexplained loss of more than 10% of body weight within 6 months preceding diagnosis.

Pathologic Classification

The clinico-pathologic characteristics for children and adults are identical:

- Nodular lymphocyte-predominant HL (nLPHL): The characteristic lymphocyte and histiocytic cells are CD20+ (B-lymphocyte marker).
 nLPHL is reminiscent of indolent non-Hodgkin lymphomas, with a lengthy time to diagnosis and time to relapse. It is relatively more common in young children, where it commonly involves a single peripheral lymph node region and spares the mediastinum.
- Lymphocyte-rich (classic) HL: Hodgkin Reed-Sternberg (R-S) cells (CD15+) are identifiable in a background predominantly of lymphocytes. Clinical behavior is similar to that of mixed-cellularity HL.
- Mixed-cellularity (classic) HL (MCHL): R-S cells (CD15+) are frequent in a background of abundant normal reactive cells (lymphocytes, plasma cells, eosinophils, histiocytes).
- Nodular sclerosis (classic) HL (NSHL): Collagenous bands divide the lymph node into nodules which often contain an R-S cell variant
 called the lacunar cell. NSHL frequently occurs in children, involving supradiaphragmatic nodes and spreading along contiguous nodal
 chains.
- Lymphocyte-depleted (classic) HL (LDHL): This subtype is rare and commonly confused with non-Hodgkin lymphoma, particularly of the anaplastic large-cell type. LDHL is often advanced at diagnosis and has a poor prognosis.

The distribution of the subtypes in younger children differs from that in adolescents and adults. Although NSHL is the most common subtype in all age groups, it is more frequent in adolescents (77%) and adults (72%) than in younger children (44%). Conversely, MCHL is more common in younger children (33%) than in adolescents (11%) or adults (17%).

Staging

The staging system is based on anatomical groups of regional lymph nodes as delineated at the 1970 Ann Arbor symposium. It was subsequently revised at the Cotswolds meeting, although not all suggestions are consistently used.

Diagnostic Evaluation

After pathologic confirmation, patients should undergo a clinical staging, beginning with a detailed history of systemic symptoms and physical examination. Laboratory studies include complete blood count and biochemical evaluation with liver function tests (including albumin). Acute-phase reactants, including erythrocyte sedimentation rate (ESR) and C-reactive protein, may be elevated at diagnosis. Patients with "B" symptoms or stage III and IV lymphoma should have a bone marrow biopsy.

Imaging studies of the neck and thorax should be performed to assess the extent of cervical and mediastinal disease. Bulky mediastinal lymphadenopathy is defined by the ratio of the mediastinal mass to the maximal measurement of the chest cavity on an upright chest radiograph; mediastinal ratios of 33% or higher are considered bulky. Computerized tomography (CT) scans are necessary to define disease involvement in the neck and chest. Distinguishing normal (or hyperplastic) thymus from nodes in young children can be challenging.

An abdominal and pelvic CT should be used for infradiaphragmatic evaluation. If CT is used, oral and intravenous contrast is required to accurately define retroperitoneal and pelvic lymph nodes. HL involving the liver or spleen is suggested by CT findings of definite areas of abnormal density representing lymphomatous deposits. If the etiology of abnormalities seen in the liver on CT is not clear, then magnetic resonance imaging (MRI) and/or positron emission tomography (PET) can be useful to aid diagnosis.

PET is increasingly recognized as the most useful functional staging modality for lymphoma. Uptake of the radioactive glucose analogue fluorodeoxyglucose (FDG) correlates with metabolic activity in tumors undergoing anaerobic glycolysis. Areas of abnormal avidity have assisted in disease delineation and been correlated with outcome when assessed after initial cycles of chemotherapy and at completion of therapy. However, there are limitations in the pediatric setting (i.e., false positives for a variety of reasons as well as false negatives in the presence of necrosis).

Prognostic Factors

As the treatment of HL has improved, factors that are associated with outcome have become more difficult to identify. However, several prognostic factors continue to influence the success and choice of therapy. Also, most data are based on reports that primarily include adults.

- The *stage of disease* persists as the most important prognostic variable. Patients with advanced-stage disease, especially stage IV, have a poorer outcome than patients with early-stage disease.
- The *bulk of disease* combines the number of disease sites and the volume of involvement at each site. Patients with *several sites* of involvement, generally defined as four or more, fare less well.
- Systemic ("B") symptoms result from cytokine secretion, reflect biologic aggressiveness, and confer a worse prognosis.
- Laboratory studies, including the ESR, hemoglobin level, and serum albumin, have been reported to predict worse outcomes. This could

reflect disease biology or bulk.

- Histologic subtype is relevant. Patients with nLPHL are biologically different as demonstrated by improved disease-free survival (DFS) and overall survival (OS); separate protocols with minimal therapy are underway for early-stage patients. Patients with LDHL fare poorly.
 Mixed reports suggest better or poorer outcome of other histologies that may be related to other prognostic factors as well.
- Age is a significant prognostic factor, with survival rates for children with HL approaching 85% to 95%. In a report from Stanford, the 5and 10-year survival rates for children with HL ≤10 years of age are 94% and 92%, respectively, compared with 93% and 86% for
 adolescents (11 to 16 years of age) and 84% and 73% for adults.
- Rapidity of response to initial therapy is an important prognostic variable. Early response to therapy was initially observed in advanced-stage HL patients treated on Pediatric Oncology Group (POG) 8725, where 93% of patients who attained a complete response (CR) after 3 cycles of chemotherapy remained disease free. This finding was also confirmed for lower-stage patients and afterwards incorporated in the latest COG front-line trials. Early CR to therapy has also been successfully incorporated into the German trials with low-risk patients who achieve CR after 2 cycles of OEPA (vincristine, etoposide, prednisone, and doxorubicin) not requiring further radiotherapy (RT). Response-based therapy is currently the paradigm on which modern pediatric trials are based.

Selection of Therapy

The desire to cure young children with minimal side effects has led to careful risk stratification, in an attempt to optimize reduced intensity and types of chemotherapy, as well as RT doses and volumes. Because of differences in the age-related developmental status of children and the gender-related sensitivity to gonadal chemotherapy toxicity, no single treatment is ideal for all children. The use of RT and chemotherapy can broaden the spectrum of potential toxicities but reduce the severity of individual toxicities. Most current approaches entail chemotherapy in conjunction with reduced RT doses. The volume of RT and the intensity and duration of chemotherapy are risk- and response-adapted and determined by prognostic factors at presentation.

Chemotherapy with mechlorethamine, vincristine, procarbazine, and prednisolone (MOPP) fell out of favor due to major toxicities, including risks of secondary acute myeloid leukemia, azoospermia in more than 90% of males treated at any age, and sterility in females, which increases with age. Doxorubicin, bleomycin, vinblastine, and dacarbazine (ABVD) became front-line chemotherapy in adults, as secondary leukemia and sterility are less common. The predominant adverse effects of ABVD are pulmonary toxicity related to bleomycin and cardiovascular toxicity secondary to doxorubicin. These side effects may be exacerbated by the addition of mediastinal or mantle irradiation. Current pediatric regimens have evolved to further limit the risks of sterility, leukemia, and cardiopulmonary toxicity.

Risk-Adapted Therapy – Favorable-Risk Disease

Favorable-risk disease is defined differently by different clinical trial groups, and even among the same groups sometimes the concept evolves over time, so that protocols are not easily comparable. For the most part, favorable-risk disease encompasses patients with localized stage I and II disease without adverse prognostic features; however this definition will vary from group to group. Some of the unfavorable features considered are "B" symptoms, extranodal extension, peripheral or mediastinal bulky disease, hilar adenopathy, and 3 or more nodal regions. Treatment typically involves 2 to 4 cycles of chemotherapy and low-dose, involved-field radiation therapy (IFRT). In some regimens, the RT dose has been reduced based on a favorable response to chemotherapy (see Appendix 1 in the original guideline document).

The German Paediatric Oncology and Haematology Society (GPOH) pioneered the use of risk- and gender-adapted therapy featuring the OEPA (vincristine, etoposide, prednisone, and doxorubicin) regimen for boys in order to limit the amount of alkylators, while girls received OPPA (vincristine, procarbazine, prednisone, and doxorubicin). The GPOH HD-95 trial investigated whether RT could be omitted in patients achieving a CR to chemotherapy. Early results (median follow-up time of 3 years) indicate a 97% event-free survival (EFS) rate for favorable-risk patients. There was no difference in outcome between favorable-risk patients treated with chemotherapy alone and those treated with combined-modality therapy. Importantly, the criteria for CR were strict (CR defined as a volume reduction of ≥95% and ≤2 mL of the initial volume or unconfirmed CR if volume reduction was ≥75% or <2 mL) so that less than 30% of the favorable-risk patients fell into this category; the cohort consisted of classical HL and nLPHL. These results were confirmed in the GPOH-HD 2002 study that excluded LPHL patients. In this study, all patients received IFRT to 19.8 Gy except those in the early-stage (IA/B and IIA without extranodal involvement) disease category who achieved a CR after induction therapy as defined in GPOH-HD 95. In regions with <75% volume reduction, a boost to approximately 30 Gy was administered, and residual masses >100 mL were boosted to approximately 35 Gy. As in the previous study, less than one-third of all early-stage patients achieved a CR by these strict criteria.

Several North American investigators have observed excellent treatment results in combined-modality trials for favorable-risk HL. Pediatric Hodgkin consortium investigators from Stanford, St. Jude, and Dana Farber reported treatment results using a nonalkylator regimen, VAMP (vinblastine, doxorubicin, methotrexate, and prednisone) for children with clinical I/II, nonbulky HL. Patients received 4 cycles of VAMP chemotherapy and response-based IFRT after 2 cycles of chemotherapy. At a median follow-up of 9.6 years, 5- and 10-year EFS rates were 92.7% and 89.4%, respectively. The Pediatric Oncology Group (POG) evaluated the feasibility of combined-modality therapy using 4 courses of

DBVE (doxorubicin, bleomycin, vincristine, and etoposide) followed by IFRT to 25.5 Gy to treat stage IA, IIA, and IIIA HL. At a median follow-up of 8.4 years, 6-year OS and EFS rates were 98% and 91%, respectively, with almost all patients (98%) achieving remission after completion of therapy. This DBVE regimen was used by the POG and the Children's Oncology Group (COG) to support reduction of chemotherapy via an early-response-based treatment algorithm. Patients received only 2 courses of ABVE if they achieved an early CR (45% of all patients) versus 4 courses of ABVE if they were slower responders. IFRT to 25.5 Gy was subsequently given to all, resulting in 5-year OS and EFS of 98% and 88%, respectively. In the COG AHOD0431 single-arm study, 287 stage IA/IIA patients who achieved a CR after 3 cycles of doxorubicin, vincristine, prednisone and cyclophosphamide (AVPC) received no further therapy. Those with a partial response (PR) received 21 Gy IFRT. The 2-year EFS rate was 80% for those achieving CR after 3 cycles of AVPC (i.e., no RT) versus 88% for patients achieving PR (and receiving IFRT) (P=0.11). The 2-year OS rate was 100%. Of the evaluable patients with fluorine-18-2-fluoro-2-deoxy-D-glucose positron emission tomography (FDG-PET) results after one cycle of chemotherapy (PET1), the 2-year EFS rates for CR patients who had a positive/equivocal PET1 versus who had a negative PET1 were 65% versus 87%, respectively (P=0.005). The 2-year EFS rates for PR patients who had a positive/equivocal PET1 versus a negative PET1 were 82% versus 96%, respectively (P=0.047). These preliminary results suggest that CT response alone is not adequate to identify patients who can be treated without RT after abbreviated chemotherapy. However, very early response as measured by PET1 may be useful for developing response-adapted therapy in favorable-risk patients.

In the Children's Cancer Group (CCG) trial, chemotherapy alone using the COPP/ABV (cyclophosphamide, vincristine, procarbazine, prednisone, doxorubicin, bleomycin, and vinblastine) hybrid regimen was compared to risk-adapted combined-modality therapy with low-dose IFRT. Patients achieving a CR to chemotherapy were eligible for randomization to receive low-dose IFRT or no further therapy. The trial was terminated early due to a significantly higher number of relapses among patients treated with chemotherapy alone. The 3-year EFS estimates were 92% for patients treated with combined-modality therapy and 87% for those treated with chemotherapy alone. The benefit of IFRT remained significant in the "as treated" analysis. Estimates of OS are not different between the randomized groups in early follow-up; however, salvage therapy after relapse is a known risk for neoplastic complications and early mortality. In the GPOH-HD 95 trial, the relapse-free survival rate was better for patients treated with RT after PR (93%) than for those without RT after CR (89%). The difference was significant for patients treated for advanced-stage but not early-stage disease. These results were confirmed in the GPOH-HD2002 trial. Patients with stage I, IIA, or IIIA HL were randomized in a POG 8625 study after a CR or PR to four courses of MOPP/ABVD to either two additional courses of alternating MOPP/ABVD or to IFRT to 25.5 Gy. At a median follow-up of 8.25 years, 8-year EFS rates were 83% for chemotherapy alone and 91% for combined-modality therapy, while 8year OS rates were 93.6% for chemotherapy alone and 96.8% for chemotherapy and RT. This study was powered to detect a 15% difference in 3-year EFS rates with 80% power in comparison to the CCG trial which was powered to detect a 6% difference in postrandomization EFS rates with 83% power. Therefore, despite the larger difference in EFS between the two arms in the POG trial compared to the CCG 5942 trial, these differences were not statistically different. In the POG study, patients with early response to therapy had a significantly better outcome, which supports the paradigm of response-based treatment (see Variant 1 above).

Risk-Adapted Therapy – Intermediate-Risk Disease

In risk-adapted treatment regimens, patients presenting with localized (stage IA, IIA) disease with unfavorable features are often grouped into an intermediate-risk category that also includes those stage IIIA disease. The GPOH-HD84 was the first study to give this group of patients less intensive therapy than the unfavorable group, but more intense therapy compared to the favorable group. Building on this concept, the GPOH-HD 2002 study reported a 5-year EFS rate of 88% with 2 cycles of OEPA for boys and OPPA for girls. This was followed by 2 cycles of COPP for girls and cyclophosphamide, vincristine, prednisone, and dacarbazine (COPDAC) for boys in order to spare fertility by reducing alkylator exposure. The COG has developed an approach of using dose density to support early-response-adapted therapy. A dose-dense regimen of doxorubicin, bleomycin, vincristine, etoposide, prednisone, and cyclophosphamide (ABVE-PC) chemotherapy was used for both intermediate and high-risk patients in the POG9425 study. This study provided encouraging clinical outcomes using response-based therapy. Patients with a rapid early response (RER) to 3 cycles of ABVE-PC received 21 Gy of regional field RT (mantle, para-aortic, or pelvis). Slow early responders (SER) received an additional 2 cycles of ABVE-PC and then received IFRT to 21 Gy. The 5-year EFS rates were 86% for RER patients and 83% for SER patients (P=0.85), and the 5-year OS rate was 95%. In the recently completed COG AHOD 0031 study, patients received 2 cycles of ABVE-PC followed by response assessment. Patients with RER received 2 additional cycles of ABVE-PC followed by a second response assessment. Those with a CR were randomized to 21 Gy IFRT or no further therapy. Patients with a RER who did not have a CR were all assigned to receive IFRT. SER patients were all randomized to either 2 additional cycles of ABVE-PC or dexamethasone, etoposide, cisplatin, and cytarabine (DECA) followed by an additional 2 cycles of ABVE-PC. All SER patients received 21 Gy IFRT after chemotherapy. Three-year EFS rates were 87.1% for RER patients versus 77.8% for SER patients (P=0.0001). The 3-year OS rate for RER patients was 98.7% versus 96.9% for SER patients (P=0.02). The 3-year EFS rate was 87.9% for RER/CR patients randomized to receive IFRT versus 85.4% for those randomized to no IFRT (P=0.07). These results suggest that early response to chemotherapy defined by early reduction (60%) in tumor size based on CT after 2 cycles can be a powerful predictor of outcome and help optimize subsequent treatment. A secondary analysis of PET response after 2 cycles of ABVE-PC demonstrated that PET may further assist with treatment optimization. Analyses of the AHOD 0031 cohort are still ongoing, including an "as treated" analysis. This will yield further information on the influence of disease characteristics, such as bulk, and treatmentrelated factors on clinical outcomes (see Variant 2 and Variant 3 above).

Risk-Adapted Therapy – Unfavorable-Risk Disease

The criteria for unfavorable clinical presentations vary, but typically include the presence of "B" symptoms, bulky lymphadenopathy, hilar lymphadenopathy, involvement of 3 or more nodal regions, extranodal extension to contiguous structures, or advanced-stage (IIIB-IV). RT for unfavorable and advanced HL is variable and protocol dependent (see Appendix 2 in the original guideline document). Although IFRT remains the standard in patients treated with combined-modality therapy, restriction of RT to areas of initial bulky disease or postchemotherapy residual disease is under investigation.

For patients with unfavorable or advanced disease, two primary treatment approaches have been used. A conventional treatment approach involves chemotherapy on a twice-monthly schedule for 6 to 8 months. An alternative strategy condenses treatment into 3 to 5 months to enhance dose intensity and reduce the risk of developing resistant disease. Dose-intensified treatment regimens may also increase the risk of acute and late side effects. A recent assessment of a dose-intense, response-based regimen using BEACOPP (bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone) in children with high-risk disease was reported by the COG. Females with a rapid response received 4 cycles of COPP/ABV following BEACOPP without IFRT, and males received 2 cycles of ABVD and IFRT. All patients with a slow response received 4 additional cycles of BEACOPP and IFRT. A high 5-year EFS rate of 94% was achieved, and the 5-year OS rate was 97%. A summary of treatment results of published trials is provided in Appendix 2 of the original guideline document, which demonstrates EFS rates ranging from 70% to 90% (see Variant 4 above).

The GPOH, building on its original experience with the OPPA/COPP regimen, showed that 6 cycles of OEPA/COPDAC together with 20-30 Gy IFRT also produced excellent results, with a 5-year EFS rate of approximately 87%.

In summary, early results have suggested that response-adapted therapy may identify favorable-risk pediatric HL patients who can be treated with chemotherapy alone without significantly reducing DFS. In the past, chemotherapy-alone regimens for advanced-stage disease used higher cumulative doses, predisposing survivors to greater risks of acute and late toxicity associated with alkylating agents, anthracyclines, and bleomycin. These protocols were designed with the hopes of avoiding toxicities due to RT, including cardiopulmonary dysfunction and solid-tumor carcinogenesis. Current protocols have carefully balanced chemotherapy agents, treatment modalities, and doses to limit long-term risks. The data suggest that children with advanced or unfavorable symptomatic or bulky disease at presentation have better outcomes using a combined-modality approach. Identification of prognostic features requiring RT to optimize disease control is a focus of many ongoing pediatric trials.

Radiotherapeutic Management

Most newly diagnosed children will be treated with risk-adapted chemotherapy alone or with combined-modality therapy, including low-dose IFRT. In the past, fully grown adolescents with favorable early-stage disease were treated with full-dose extended-field radiation therapy (EFRT) using techniques that are standard for adults. However, this approach has been abandoned due to concerns of cardiac toxicity and second cancers.

Radiation fields must be meticulously and judiciously designed to maximize disease control and minimize normal tissue damage. Field definition depends on the anatomy of the region, including node distribution and patterns of disease extension. The traditional definitions of lymph node regions can be helpful but are not necessarily sufficient. As a result, field definitions are often protocol-specific.

Efforts to exclude unnecessary normal tissues (e.g., breast tissue) are always important in a child with isolated mediastinal disease and no axillary involvement. Involved supradiaphragmatic fields can be simulated with the arms above the head, or down with hands on the hips. The former pulls the axillary lymph nodes away from the lungs, allowing greater lung shielding; however, axillary lymph nodes then move into the vicinity of the humeral heads, which should be blocked in growing children. Breast tissue should be excluded or positioned under the lung/axillary blocking. When the decision is made to include some or all of a critical normal organ in the radiation field (e.g., liver, kidney, or heart), normal tissue constraints are critical, particularly when boosting to doses >20 Gy. One should also take into account the chemotherapy protocol used.

While CT-based planning and anterior-posterior opposed parallel pair beam arrangements remain common for radiation delivery in pediatric HL, three-dimensional conformal radiation therapy (3DCRT) using nonopposed beams, intensity-modulated radiation therapy (IMRT), or proton therapy may be considered in situations where more conformal techniques would reduce dose to surrounding normal critical structures. This is sometimes the case when treating the thorax to spare dose to the heart, lungs, and developing breast tissue, or when treating the abdomen and pelvis to minimize dose to the highly radiosensitive reproductive organs. Although data are accumulating in regard to the efficacy of IMRT and the decrease in median dose to normal surrounding tissues, some uncertainty exists about the potential for increased late effects from IMRT, particularly secondary malignancy, since IMRT results in a lower dose to a larger volume compared to conventional techniques. Therefore, keeping dosimetric parameters such as the V5 to breast and lung tissue to as low as reasonably achievable may be relevant as well for HL patients. Data also suggest that use of a mean lung dose constraint <15 Gy results in low rates of radiation pneumonitis. Proton therapy is currently being

investigated and may further decrease the mean dose to normal surrounding tissue compared with IMRT or 3DCRT without increasing the volume of normal tissue receiving lower dose radiation. Although the efficacy of these more conformal techniques have been shown in other disease sites, the benefits of lower dose to critical organs are unlikely to be fully appreciated for a couple of decades.

Because patients with early-stage HL treated with chemotherapy alone most frequently relapse in the initially involved lymph node(s), efforts have been made to reduce treatment fields to include only the initially involved lymph node(s) and exclude surrounding normal tissues. The EORTC-GELA introduced the concept of involved-node radiation therapy (INRT), which uses all available clinical information, including pre- and post-chemotherapy imaging with CT with the patient in the treatment position and FDG-PET scan to define the treatment field. The pre-chemotherapy PET/CT scan should be performed in the same treatment position as when the radiation will be delivered for accurate definition of the INRT field. Controversy still exists regarding the optimum margins for INRT, but initial clinical data are emerging. A group of researchers reviewed clinical outcomes of patients with limited-stage HL treated with EFRT, IFRT, and INRT and found no marginal recurrences or locoregional failures with INRT. However, the INRT fields in this study employed margins \leq 5 cm and used conventional treatment planning. These fields were thus significantly larger than the margins prescribed for INRT by the EORTC-GELA and German Hodgkin Study Group (GHSG) guidelines. INRT requires that all available clinical information be used to appropriately reduce treatment field size without compromising the excellent clinical outcomes attainable with standard therapy.

Summary Recommendations for Primary Disease

Optimal treatment planning involves a multidisciplinary approach beginning at diagnosis, with the pediatric and radiation oncologist meeting to review staging studies with a radiologist following examination of the patient. The treatment approach should consider patient factors such as age and gender that may enhance the risk of complications, as well as disease factors (e.g., presence of "B" symptoms, bulky lymphadenopathy, and stage). Recommended treatment approaches for favorable localized, intermediate, and advanced unfavorable disease are summarized in the Table below (see Variant 5 above).

Table. Recommendations for Treatment Approach in Pediatric Hodgkin Lymphoma

Clinical Presentation	Stage	Recommended treatment approach
Low Risk: Localized disease involving <3-4 nodal regions in absence of "B" symptoms, bulk, or extranodal extension	IA, IIA	Recommended therapy: 2-4 cycles non-cross-resistant chemotherapy (OEPA, VAMP, COPP-ABV, AVPC). Response-based low-dose, IFRT (1500 cGy-2550 cGy). Other considerations: Consider use of IFRT based on early response to chemotherapy. If CR after 2 cycles of OEPA, no need for IFRT AV-PC without RT for RER.
Intermediate: Localized disease involving ≥3-4 nodal regions in presence of bulky lymphadenopathy (mediastinal ratio ≥33%; lymph node mass ≥6-10 cm), extranodal extension.	IA, IIA, IB*, IIIA	Recommended therapy: 3-6 cycles compacted, dose-intensive, non-cross-resistant chemotherapy (OEPA/COPP, ABVE-PC) plus low-dose, IFRT (1500 cGy-2550 cGy). Other considerations: Early response to therapy may be considered in determining need for radiation in those achieving CR.
High Risk: Stage II patients with constitutional symptoms of fever or weight loss or any patient with advanced stage	IIB*, IIIB, IV	Recommended therapy: 4-6 cycles compacted, dose-intensive cycles of non-cross-resistant chemotherapy (COPP/OEPA, ABVE-PC) plus low-dose IFRT (1500 cGy-2550 cGy). Other considerations: 8 cycles non-cross-resistant chemotherapy alone (BEACOPP) for high risk, poor early response.

^{*}Stage IIB patients have been variably treated as intermediate or unfavorable risk. Some studies use associated factors, e.g., weight loss, bulk disease, extranodal extension, for further risk stratification.

Note: Abbreviations used in the tables are listed at the end of the "Major Recommendations" field.

Nodular Lymphocyte-Predominant Hodgkin Lymphoma

Patients with nLPHL typically present with early-stage disease, peripheral lymph node (cervical, axillary, or inguinal) involvement, and a striking male predominance. The disease is considered indolent with a favorable prognosis characterized by late recurrences. The rare deaths are related to secondary malignancies, cardiopulmonary treatment toxicity, or transformation to aggressive B-cell lymphoma. Historically, patients with nLPHL have been treated on protocols for patients with classical HL; however, there are several reports in the adult and pediatric literature suggesting that

they can be cured with less aggressive therapies. In adults, radiotherapy-only approaches are favored with reasonable outcomes; however, the required doses of 30 to 40 Gy would be unacceptable in a pediatric setting except in very rare clinical circumstances.

In pediatrics, only small retrospective series have been published reviewing therapy (resection, radiotherapy, or chemotherapy alone, and combined-modality) and outcome of these patients, most of whom do well regardless of the modality chosen; however, toxicity remains the main concern for more involved treatment approaches. The largest retrospective study evaluating resection alone in children looked at the outcome of 58 children from several European countries. This study proved that a substantial proportion of patients with limited-stage nLPHL and completely resected disease could achieve a long-term remission without any further therapy. Based on these findings, the Euro-Net is conducting a clinical trial to prospectively validate this "watch and wait" approach for children with completely resected disease and low-dose chemotherapy for patients with residual disease after resection. The COG recently released the data of their AHOD03P1 study on 52 stage IA patients who were observed after surgery only. Nine relapses were observed among these patients and all were retrieved successfully with 3 cycles of AVPC (doxorubicin, vincristine, prednisone and cyclophosphamide) and no radiotherapy. The current 2-year EFS estimate among these patients is 80.3% (95% confidence interval [CI]: 65.3%-89.3%).

Refractory and Relapsed Disease

HL may still be cured if initial treatment programs fail. Relapse occurs most often within 4 years, but late relapse is not rare, especially in nLPHL patients who can fail 10 years after initial diagnosis. The spectrum of treatment options include standard-dose chemotherapy (with or without RT), RT alone, or high-dose chemotherapy (with or without RT), followed by stem-cell support, clinical trials, or palliative therapy. RT may also be used pre- or post-transplant depending on the clinical scenario.

Factors that independently predict a more favorable outcome include the site of relapse (nodal better than extranodal), stage at relapse (early better than advanced), histology, and response to first-line salvage chemotherapy. The selection of the most appropriate salvage regimen is based on whether a complete remission was achieved, the durability of the remission, the extent of disease at relapse, and the intensity of the frontline therapy given. Three- to 5-year survival probabilities of 25% to 80% have been reported (primarily in adults) following treatment with high-dose chemotherapy and hematopoietic stem cell rescue. Data specific to children with recurrent HL are limited. One study reported probabilities of relapse at 2 and 5 years of 36% (+/-5%) and 44% (+/-6%), respectively. Progression-free survival (PFS) rates were 40% (+/-6%) and 30% (+/-6%) and OS rates were 54% (+/-6%) and 45% (+/-6%) at 2 and 5 years, respectively.

For higher risk relapses, a combination of ifosfamide and vinorelbine for pediatric patients in first relapse was studied by the COG (AHOD00P1). This regimen showed a very good overall response rate (CR/PR) of 78% and achieved good stem cell mobilization for future autologous stem cell transplant. The goal in this group of patients is to proceed to autologous stem cell transplant once remission is achieved, since studies have shown that patients undergoing autologous stem cell transplant with active disease have a worse outcome. For patients who relapse after transplant or are upfront refractory, a combination of gemcitabine and vinorelbine was also studied by this group. It resulted in an overall response rate of 76%. Patients who relapse after an autologous stem cell transplant are often considered for an allogeneic stem cell transplant.

A group of researchers found an increased risk of relapse and lower PFS beyond 9 months when pediatric patients received reduced intensity conditioning compared to a myeloabalative conditioning regimen prior to undergoing allogeneic stem cell transplant. OS, however, was not different between the two groups.

Newer drugs promise great efficacy with less toxicity. Targeted therapy with brentuximab vedotin, an antibody-drug conjugate that targets CD30, has shown excellent results in early clinical trials. Pediatric trials are underway to assess its efficacy and toxicity, and discussions about incorporating it into large clinical trials are under way. Furthermore, histone deacetylase (HDAC) inhibitors like panobinostat are being investigated, as well as mammalian target of rapamycin (mTOR) inhibitors.

Summary

- In an effort to cure children with minimal side effects, most current treatment approaches to pediatric HL entail combined-modality therapy with reduced dose radiation.
- Prognostic factors at presentation are used to risk stratify patients and determine treatment approach.
- A number of combined-modality therapy protocols exist, and the decision as to how to incorporate RT should be made within the context of the protocol followed.
- For favorable-risk disease, 2 to 4 cycles of non-cross-resistant chemotherapy are recommended, followed by response-based low-dose (e.g., 15-25 Gy) IFRT depending on the protocol used.
- For intermediate-risk disease, 3 to 6 cycles of compacted, dose-intensive chemotherapy are recommended. Early response to therapy may be considered in determining the need for radiation in those achieving CR.
- For high-risk disease, 4 to 6 compacted, dose-intensive cycles of chemotherapy in addition to low-dose (e.g., 15-25 Gy) IFRT to involved sites of disease are recommended.

- Involved-node radiotherapy (INRT) in the pediatric population remains investigational.
- Radiotherapy-alone approaches for pediatric nLPHL are discouraged due to the higher doses (e.g., 30-40 Gy) required that may affect skeletal maturity. The approach of resection alone for limited-stage disease appears promising, although it is still under investigation.
 Chemotherapy-alone approaches remain standard.
- Advanced radiation techniques, such as IMRT and proton therapy, may be considered depending on the clinical scenario and if an
 improvement in the therapeutic ratio is expected.

Abbreviations

- ABV, adriamycin (doxorubicin), bleomycin, and vinblastine
- ABVD, adriamycin (doxorubicin), bleomycin, vinblastine, and dacarbazine
- ABVE-PC, adriamycin (doxorubicin), bleomycin, vincristine, etoposide, prednisone, and cyclophosphamide
- BEACOPP, bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone
- COG, Children's Oncology Group
- COPP, cyclophosphamide, oncovin (vincristine), prednisone, and procarbazine
- COPDac, cyclophosphamide, oncovin (vincristine), prednisone, and dacarbazine
- CR, complete remission
- CS, clinical stage
- DBVE, doxorubicin, bleomycin, vincristine, and etoposide
- IFRT, involved-field radiation therapy
- IMRT, intensity-modulated radiation therapy
- MCHL, mixed-cellularity Hodgkin lymphoma
- nLPHL, nodular lymphocyte-predominant Hodgkin lymphoma
- NSHL, nodular sclerosis Hodgkin lymphoma
- OEPA, oncovin (vincristine), etoposide, prednisone, and adriamycin (doxorubicin)
- OPPA, oncovin (vincristine), procarbazine, prednisone, and adriamycin (doxorubicin)
- RER, rapid early response
- RT, radiotherapy
- VAMP, vinblastine, adriamycin (doxorubicin), methotrexate, and prednisone

Clinical Algorithm(s)

Algorithms were not developed from criteria guidelines.

Scope

Disease/Condition(s)

Hodgkin lymphoma

Guideline Category

Management

Risk Assessment

Treatment

Clinical Specialty

Oncology

Radiation Oncology	
Radiology	
ntended Users	

Health Plans

Pediatrics

Hospitals

Managed Care Organizations

Physicians

Utilization Management

Guideline Objective(s)

To evaluate the appropriateness of radiologic treatment procedures and chemotherapy regimens for pediatric patients with Hodgkin lymphoma

Target Population

Children with Hodgkin lymphoma

Interventions and Practices Considered

- 1. Risk-adapted chemotherapy with or without radiation therapy, including consideration of type and duration
- 2. Radiation therapy alone (not generally recommended)
- 3. Surgery alone

Major Outcomes Considered

- Overall survival rate
- Disease-free, relapse-free, and event-free survival rate
- Risk of relapse
- Toxicity of treatment

Methodology

Methods Used to Collect/Select the Evidence

Searches of Electronic Databases

Description of Methods Used to Collect/Select the Evidence

Literature Search Procedure

The Medline literature search is based on keywords provided by the topic author. The two general classes of keywords are those related to the condition (e.g., ankle pain, fever) and those that describe the diagnostic or therapeutic intervention of interest (e.g., mammography, MRI).

The search terms and parameters are manipulated to produce the most relevant, current evidence to address the American College of Radiology Appropriateness Criteria (ACR AC) topic being reviewed or developed. Combining the clinical conditions and diagnostic modalities or therapeutic procedures narrows the search to be relevant to the topic. Exploding the term "diagnostic imaging" captures relevant results for diagnostic topics.

The following criteria/limits are used in the searches:

- 1. Articles that have abstracts available and are concerned with humans.
- 2. Restrict the search to the year prior to the last topic update or in some cases the author of the topic may specify which year range to use in the search. For new topics, the year range is restricted to the last 5 years unless the topic author provides other instructions.
- 3. May restrict the search to Adults only or Pediatrics only.
- 4. Articles consisting of only summaries or case reports are often excluded from final results.

The search strategy may be revised to improve the output as needed.

Number of Source Documents

The total number of source documents identified as the result of the literature search is not known.

Methods Used to Assess the Quality and Strength of the Evidence

Weighting According to a Rating Scheme (Scheme Given)

Rating Scheme for the Strength of the Evidence

Strength of Evidence Key

- Category 1 The conclusions of the study are valid and strongly supported by study design, analysis and results.
- Category 2 The conclusions of the study are likely valid, but study design does not permit certainty.
- Category 3 The conclusions of the study may be valid but the evidence supporting the conclusions is inconclusive or equivocal.
- Category 4 The conclusions of the study may not be valid because the evidence may not be reliable given the study design or analysis.

Methods Used to Analyze the Evidence

Systematic Review with Evidence Tables

Description of the Methods Used to Analyze the Evidence

The topic author drafts or revises the narrative text summarizing the evidence found in the literature. American College of Radiology (ACR) staff draft an evidence table based on the analysis of the selected literature. These tables rate the strength of the evidence for all articles included in the narrative text.

The expert panel reviews the narrative text, evidence table, and the supporting literature for each of the topic-variant combinations and assigns an appropriateness rating for each procedure listed in the table. Each individual panel member forms his/her own opinion based on his/her interpretation of the available evidence.

More information about the evidence table development process can be found in the ACR Appropriateness Criteria® Evidence Table Development document (see the "Availability of Companion Documents" field).

Methods Used to Formulate the Recommendations

Expert Consensus (Delphi)

Description of Methods Used to Formulate the Recommendations

Modified Delphi Technique

The appropriateness ratings for each of the procedures included in the Appropriateness Criteria topics are determined using a modified Delphi methodology. A series of surveys are conducted to elicit each panelist's expert interpretation of the evidence, based on the available data, regarding the appropriateness of an imaging or therapeutic procedure for a specific clinical scenario. American College of Radiology (ACR) staff distributes surveys to the panelists along with the evidence table and narrative. Each panelist interprets the available evidence and rates each procedure. The surveys are completed by panelists without consulting other panelists. The ratings are a scale between 1 and 9, which is further divided into three categories: 1, 2, or 3 is defined as "usually not appropriate"; 4, 5, or 6 is defined as "may be appropriate"; and 7, 8, or 9 is defined as "usually appropriate." Each panel member assigns one rating for each procedure per survey round. The surveys are collected and the results are tabulated, de-identified and redistributed after each round. A maximum of three rounds are conducted. The modified Delphi technique enables each panelist to express individual interpretations of the evidence and his or her expert opinion without excessive bias from fellow panelists in a simple, standardized and economical process.

Consensus among the panel members must be achieved to determine the final rating for each procedure. Consensus is defined as eighty percent (80%) agreement within a rating category. The final rating is determined by the median of all the ratings once consensus has been reached. Up to three rating rounds are conducted to achieve consensus.

If consensus is not reached, the panel is convened by conference call. The strengths and weaknesses of each imaging procedure that has not reached consensus are discussed and a final rating is proposed. If the panelists on the call agree, the rating is accepted as the panel's consensus. The document is circulated to all the panelists to make the final determination. If consensus cannot be reached on the call or when the document is circulated, "No consensus" appears in the rating column and the reasons for this decision are added to the comment sections.

Rating Scheme for the Strength of the Recommendations

Not applicable

Cost Analysis

A formal cost analysis was not performed and published cost analyses were not reviewed.

Method of Guideline Validation

Internal Peer Review

Description of Method of Guideline Validation

Criteria developed by the Expert Panels are reviewed by the American College of Radiology (ACR) Committee on Appropriateness Criteria.

Evidence Supporting the Recommendations

Type of Evidence Supporting the Recommendations

The recommendations are based on analysis of the current literature and expert panel consensus.

Benefits/Harms of Implementing the Guideline Recommendations

Potential Benefits

Potential Harms

- The predominant adverse effects of ABVD (doxorubicin, bleomycin, vinblastine, and dacarbazine) are pulmonary toxicity related to bleomycin and cardiovascular toxicity secondary to doxorubicin. These side effects may be exacerbated by the addition of mediastinal or mantle irradiation. Current pediatric regimens have evolved to further limit the risks of sterility, leukemia, and cardiopulmonary toxicity.
- Dose-intensified chemotherapy regimens may increase the risk of acute and late side effects.
- Some uncertainty exists about the potential for increased late effects from intensity-modulated radiation therapy (IMRT), particularly
 secondary malignancy, since IMRT results in a lower dose to a larger volume compared to conventional techniques. Data also suggest that
 use of a mean lung dose constraint <15 Gy results in low rates of radiation pneumonitis.
- Because of differences in the age-related developmental status of children, and the gender-related sensitivity to gonadal chemotherapy
 toxicity, no single treatment is ideal for all children. The use of radiation and chemotherapy can broaden the spectrum of potential toxicities,
 but reduce the severity of individual toxicities.
- Positron emission tomography has limitations in the pediatric setting (i.e., false positives for a variety of reasons as well as false negatives in the presence of necrosis).

Qualifying Statements

Qualifying Statements

The American College of Radiology (ACR) Committee on Appropriateness Criteria and its expert panels have developed criteria for determining appropriate imaging examinations for diagnosis and treatment of specified medical condition(s). These criteria are intended to guide radiologists, radiation oncologists, and referring physicians in making decisions regarding radiologic imaging and treatment. Generally, the complexity and severity of a patient's clinical condition should dictate the selection of appropriate imaging procedures or treatments. Only those examinations generally used for evaluation of the patient's condition are ranked. Other imaging studies necessary to evaluate other co-existent diseases or other medical consequences of this condition are not considered in this document. The availability of equipment or personnel may influence the selection of appropriate imaging procedures or treatments. Imaging techniques classified as investigational by the U.S. Food and Drug Administration (FDA) have not been considered in developing these criteria; however, study of new equipment and applications should be encouraged. The ultimate decision regarding the appropriateness of any specific radiologic examination or treatment must be made by the referring physician and radiologist in light of all the circumstances presented in an individual examination.

Implementation of the Guideline

Description of Implementation Strategy

An implementation strategy was not provided.

Institute of Medicine (IOM) National Healthcare Quality Report Categories

IOM Care Need

Getting Better

Living with Illness

IOM Domain

Effectiveness

Identifying Information and Availability

Bibliographic Source(s)

Terezakis SA, Metzger ML, Constine LS, Hodgson DC, Schwartz CL, Advani R, Flowers CR, Hoppe BS, Ng A, Roberts KB, Shapiro R, Wilder RB, Yunes MJ, Expert Panel on Radiation Oncology-Lymphoma. ACR Appropriateness Criteria® pediatric Hodgkin lymphoma. [online publication]. Reston (VA): American College of Radiology (ACR); 2012. 16 p. [54 references]

Adaptation

Not applicable: The guideline was not adapted from another source.

Date Released

1999 (revised 2012)

Guideline Developer(s)

American College of Radiology - Medical Specialty Society

Source(s) of Funding

The American College of Radiology (ACR) provided the funding and the resources for these ACR Appropriateness Criteria®.

Guideline Committee

Committee on Appropriateness Criteria, Expert Panel on Radiation Oncology-Lymphoma

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Financial Disclosures/Conflicts of Interest

Not stated

Guideline Status

This is the current release of the guideline.

This guideline updates a previous version: Constine LS, Hoppe RT, Colman M, Deming RL, Mendenhall NP, Morris DE, Ng A, Wolkov HB,

Guideline Availability
Electronic copies: Available from the American College of Radiology (ACR) Web site
Print copies: Available from the American College of Radiology, 1891 Preston White Drive, Reston, VA 20191. Telephone: (703) 648-8900.
Availability of Companion Documents
The following is available:
 ACR Appropriateness Criteria®. Overview. Reston (VA): American College of Radiology; 2 p. Electronic copies: Available in Portable Document Format (PDF) from the American College of Radiology (ACR) Web site ACR Appropriateness Criteria®. Literature search process. Reston (VA): American College of Radiology; 1 p. Electronic copies: Available in PDF from the ACR Web site ACR Appropriateness Criteria®. Evidence table development – diagnostic studies. Reston (VA): American College of Radiology; 2013 Nov. 3 p. Electronic copies: Available in PDF from the ACR Web site ACR Appropriateness Criteria®. Evidence table development – therapeutic studies. Reston (VA): American College of Radiology; 2013 Nov. 4 p. Electronic copies: Available in PDF from the ACR Web site ACR Appropriateness Criteria® pediatric Hodgkin lymphoma. Evidence table. Reston (VA): American College of Radiology; 2012. 29 p. Electronic copies: Available from the ACR Web site
Patient Resources
None available
NGC Status
This NGC summary was completed by ECRI on August 28, 2006. This summary was updated by ECRI Institute on February 26, 2008 following the U.S. Food and Drug Administration advisory/voluntary market withdrawal of the liquid formulation of Leukine (sargramostim). This summary was updated by ECRI Institute on May 22, 2013.
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